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A series of 9 quinolines and 18 styrylquinolines was evaluated for the drugs' in vitro antileishmanial activities and cytotoxicities. The 7-aroylstyrylquinoline scaffold appeared to be the most promising one, with the most interesting compound, no. 35, exhibiting a 50% inhibitory concentration (IC $_{50}$) of 1.2 μ M and a selectivity index value of 121.5. Compound 35 was 10-fold and 8-fold more active than miltefosine and sitamaquine, the reference compounds, with selectivity indexes 607-fold and 60-fold higher, respectively.

Leishmaniasis is a family of parasitic diseases that affect about 12 million people in tropical and subtropical areas in the form of three clinical expressions: visceral leishmaniasis, which is fatal in the absence of treatment; muco-cutaneous leishmaniasis; and cutaneous leishmaniasis, which is often selfcuring. Classical drugs such as antimonials (Pentostam and Glucantime) are toxic, and drug resistance is increasing dangerously in the field (3). A liposomal amphotericin B formulation (AmBisome) less toxic than amphotericin B deoxycholate is gradually becoming the first-line therapy, especially in immunocompromised patients, but this drug must be administered by a parenteral route (11). Miltefosine (Impavido) was the first drug registered against visceral leishmaniasis in the last decade; however, its toxicity and the appearance of drug resistance justify the search for new chemical series in order to find an orally safe and active drug (8).

Quinolines substituted at the 2-position have shown *in vivo* activities against *Leishmania donovani*, and many compounds have been synthesized over the last decade (14). The Drug for Neglected Diseases Initiative (DNDi) has been considering this series for evaluation in preclinical development for about a year and a half. However, although promising, the series still requires improvements, and here we report the *in vitro* antileishmanial evaluation of new quinoline derivatives, including 2-[2-aryl(ethenyl)]-substituted quinoline (2-styrylquinolines) bearing additional aroyl/acyl groups at the C-7 position. In addition, some compounds within this series were recently shown to display substantial antiviral activity in HIV-infected cells (13, 22).

The synthesis of most of the compounds has been previously reported. Briefly, Kolbe carbonation of 8-hydroxyquinaldine afforded the pivotal hydroxyacid compound 1 (9), which was further elaborated into the 5-iodoquinaldine compound 12 and

amide compound 17 (13). Similarly, bromination of the C-5 position and protection of the salicylic moiety provided 5-bromoquinaldine compound 2, which was engaged in a modified Suzuki cross-coupling reaction to give 5-arylated derivatives 14 to 16 (19). Styrylquinoline compounds 19 to 27 were prepared from the corresponding quinaldine compound 3 by Perkin-type condensation in refluxing acetic anhydride, followed by hydrolysis in a pyridine-water mixture (10, 16, 21, 22). Finally, the 7-aroyl-stryryquinoline derivatives 28 to 35 were obtained via the 7-bromostyrylquinoline compound 5 according to a three-step sequence involving lithiation followed by condensation with the required aldehyde, manganese dioxide oxidation, and deprotection (15) (Fig. 1).

The antileishmanial evaluation of these compounds was then performed on Leishmania donovani amastigotes by using the luciferase-transfected Leishmania donovani (strain MHOM/ IN/80/Dd₈) promastigotes maintained in the laboratory of the Division of Parasitology, Central Drug Research Institute, Lucknow, India, since 2005 as described by Sunduru et al. (20). In order to assess the activity of compounds against the amastigote stage of the parasite, the mouse macrophage cell line J-774A.1, infected with promastigotes expressing the luciferase firefly reporter gene, was used. Cells were seeded in a 96-well plate at a density of 4×10^4 cells per ml in a final volume of 100 μl in RPMI 1640 containing 10% fetal calf serum, and the plates were incubated at 37°C in a CO₂ incubator. After 24 h, the medium was replaced with fresh medium containing stationary-phase promastigotes (4 \times 10⁵/100 μ l/well). Promastigotes were engulfed by the macrophage and transformed there into amastigotes. The test compounds were added at 2-fold dilutions in up to 7 points in fresh complete medium starting from a $100 \mu M$ concentration, and the plates were incubated at 37°C in a CO₂ incubator for 72 h. After incubation, the drugcontaining medium was decanted and 50 µl phosphate-buffered saline (PBS) was added in each well and mixed with an equal volume of Steady-Glo luciferase assay substrate dissolved in Steady-Glo luciferase assay buffer. After gentle shaking for 1 to 2 min, the readings were recorded in a luminometer (1, 4, 17). The values were expressed as relative luminescence

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1778 LOISEAU ET AL. Antimicrob. Agents Chemother.

12,17

$$HO_2C$$
 OH
 OHC
 OH

FIG. 1. General synthetic scheme for the quinoline and styrylquinoline derivatives evaluated in this study.

units (RLU). Data were transformed into a graphic program (Excel). The 50% inhibitory concentration (IC_{50}) for antileishmanial activity was calculated by nonlinear regression analysis of the concentration-response curve by using the four-parameter Hill equations. The *in vitro Leishmania donovani* intramacrophage amastigote system used to evaluate the antileishmanial activity of the compounds was the most relevant one, since it takes into account the pharmacokinetics barriers that a compound has to overcome before entering the parasite.

KB cells were used to evaluate the cytotoxicity of the compounds to mammalian cells, which allowed us to determine an *in vitro* selectivity index. The cell viability was determined with the 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (MTT) assay (12). Exponentially growing KB cells at a density of 1×10^5 cells per ml in a 100- μ l final volume were incubated in a 96-well plate with test drugs for 72 h. The test compounds were added at 3-fold dilutions for up to 7 points in complete medium starting from a 400 μ M concentration and were incubated at 37°C in a humidified mixture of CO_2 and 95% air in an incubator. Podophyllotoxin was used as a reference drug, and control wells containing dimethyl sulfoxide (DMSO) without drugs were also included in the experiment. Stock solutions of compounds were initially dissolved in DMSO and further diluted with fresh complete me-

dium. After incubation, 25 μ l of MTT reagent (5 mg/ml) in PBS medium, followed by syringe filtration, was added to each well and incubated at 37°C for 2 h. At the end of the incubation period, the supernatants were removed by inverting the plate completely without disturbing the cell layer, and 150 μ l of pure DMSO was added to each well. After 15 min of shaking, the readings were recorded as absorbance at 544 nm on a microplate reader. The cytotoxic effects were expressed as 50% lethal dose (i.e., as the concentration of a compound which provoked a 50% reduction in cell viability compared to cells in culture medium alone). Fifty percent cytotoxic concentration (CC₅₀) values were estimated as previously described (5, 12). The selectivity index (SI) for each compound was calculated as the ratio between cytotoxicity (CC₅₀) and activity (IC₅₀) against *Leishmania* amastigotes.

Among the simple quinolines (Table 1) and the styrylquinoline derivatives (Table 2), three compounds exhibited an IC₅₀ for parasites of less than 10 μ M (compounds 17, 18, and 20). The most interesting compound was compound 18, with an IC₅₀ for L. donovani intramacrophage amastigotes of 4.1 μM and a selectivity index of 8.3. A clear-cut structure-activity relationship showed that the introduction of a carboxyl group at any position was responsible for both a dramatic decrease in the antileishmanial activity and a decrease in the cytotoxicity decrease. This observation was confirmed when two carboxyl groups were introduced into the same molecule, resulting in no activity and no cytotoxicity (compound 27). These results could be ascribed to an excessive hydrophilicity limiting the drugparasite membrane interactions or a reaction between the carboxyl group with some compounds of the culture medium preventing the entry of the compound into the parasite.

Among the 7-aroylstyrylquinolines (Table 3), the most interesting compound was compound 35, which exhibited an IC $_{50}$ of 1.2 μ M and a selectivity index of 121.5. Compound 35 was 10-fold and 8-fold more active than miltefosine and sitamaquine, the reference compounds, with selectivity indexes 607-fold and 60-fold higher, respectively.

Compound 34 had an IC₅₀ of $2.1~\mu M$ and a selectivity index of 27.3. These compounds exhibited the best selectivity indexes in their series, despite the presence of a nitro group. The presence of the nitro group at the meta position greatly in-

TABLE 1. In vitro antileishmanial activity and cytotoxicity results for compounds 6 and 10 to 17

$$R_2$$
 R_3
 R_4

Compound	R_1	R ₂	R ₃	R_4	IC ₅₀ (μM)	CC ₅₀ (μM)	SI (CC ₅₀ / IC ₅₀ ratio) ^a
6	Н	Br	ОН	CH ₃	16.1	75.4	4.6
10	Н	CO ₂ H	OH	Н	>100	335	< 3.3
11	H	Н	CO_2H	CH_3	>100	303.5	< 3.0
12	I	CO ₂ H	OH	CH_3	>100	331.5	< 3.3
13	H	CHO	OH	CH_3	>100	181.9	< 1.8
14	Ph	CO ₂ H	OH	CH_3	>100	329.9	< 3.2
15	4-MePh	CO ₂ H	OH	CH_3	>100	169.7	< 1.6
16	3-Quinolinyl	CO ₂ H	OH	CH_3	>100	>400	
17	Н	p-FPhCH ₂ NHCO	OH	CH_3	5.9	32.9	5.5

^a The selectivity index (SI) is defined as the ratio of CC₅₀ on KB cells to IC₅₀ on L. donovani intramacrophage amastigotes.

TABLE 2. In vitro antileishmanial activity and cytotoxicity results for styrylquinolines 18 to 27

$$R_2$$
 R_3
 R_4
 R_5

Compound	R_1	R_2	R ₃	R_4	R_5	R_6	IC ₅₀ (μM)	CC ₅₀ (μM)	SI (CC ₅₀ / IC ₅₀ ratio) ^a
18	Н	CN	ОН	Н	ОН	ОН	4.1	34.5	8.3
19	Н	Н	NO_2	Н	OAc	OAc	21.3	39.7	1.8
20	Н	Н	OH	Н	Н	Н	6.9	4.0	0.6
21	H	H	OAc	Н	OAc	OAc	13.5	9.8	0.7
22	H	CO ₂ H	OH	Н	NH_2	H	>100	35.6	< 0.3
23	H	CO_2H	OH	Н	SO ₂ CH ₃	H	>100	263.3	< 2.6
24	H	CO_2H	OH	Н	F	F	>100	193.8	< 1.9
25	H	CO_2H	OH	Н	Cl	OH	79.6	299.8	3.7
26	H	CO_2H	OH	OH	Н	OH	>100	>400	
27	CO_2H	CO_2^2H	OH	Н	OH	OH	>100	>400	

^a The selectivity index (SI) is defined as the ratio of CC₅₀ on KB cells to IC₅₀ on L. donovani intramacrophage amastigotes.

creased the selectivity index, as evidenced by the much lower selectivity index of the parent compound, no. 28.

Recent work has confirmed the interesting antileishmanial properties of other quinoline series (2, 6, 18). In addition, quinolines have recently been found to inhibit leishmanial GDP-mannose-pyrophosphorylase, an enzyme system producing a range of mannose-rich glycoconjugates that are essential for parasite survival and virulence (7). This potential for selective action against a *Leishmania*-specific target makes quinolines a promising series of antileishmanial drugs. Moreover, we have tried to select quinoline-resistant *L. donovani* promastigotes in the lab by *in vitro* drug pressure and have only obtained a slight decrease in sensitivity since the IC₅₀s were no more than twice those of the wild-type line (data not shown). This encouraging result is an additive justification for further studies of 2-substituted quinolines.

In conclusion, compound 35, due to its high in vitro anti-

TABLE 3. *In vitro* antileishmanial activity and cytotoxicity results for 7-aroylstyrylquinoline compounds 28 to 35

Compound	R_1	IC ₅₀ (μM)	CC ₅₀ (µM)	SI (CC ₅₀ / IC ₅₀ ratio) ^a
28	Ph	2.5	36.5	14.6
29	1-Naphthyl	20.0	1.1	0.06
30	PhCH ₂ CH ₂	3.9	10.3	2.6
31	3,4-DiFPh	2.5	16.5	6.5
32	2-Pyridyl	58.7	260.5	4.4
33	3-Pyridyl	17.5	9.0	0.5
34	$2-NO_2Ph$	2.1	57.4	27.3
35	$3-NO_{2}Ph$	1.2	145.8	121.5
Sitamaquine	-	9.7	19.5	2.0
Miltefosine		13.4	3.2	0.2

 $[^]a$ The selectivity index (SI) is defined as the ratio of CC₅₀ on KB cells to IC₅₀ on L. donovani intramacrophage amastigotes.

leishmanial activity and low toxicity, is the most interesting compound to emerge from more than 150 derivatives of 2-substituted quinolines that have now been synthesized and evaluated. It has now been selected as a candidate for evaluation *in vivo* with *L. donovani* mouse or hamster models via the DNDi pipeline.

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1780 LOISEAU ET AL. Antimicrob. Agents Chemother.

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